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Unique Gold Nanoparticle Aggregates as a Highly Active SERS Substrate

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ABSTRACT. A unique gold nanoparticle aggregate (GNA) system has been shown to be an excellent

substrate for surface-enhanced Raman scattering (SERS) applications. Rhodamine 6G (R6G), a

common molecule used for testing SERS activity on silver, but generally difficult to detect on gold

substrates, has been found to readily bind to the GNA and exhibit strong SERS activity due to the

unique surface chemistry afforded by sulfur species on the surface. This GNA system has yielded a

large SERS enhancement of 10⁷-10⁹ in bulk solution for R6G, on par with or greater than any previously

reported gold SERS substrate. SERS activity has also been successfully demonstrated for several

biological molecules including adenine, L-cysteine, L-lysine, and L-histidine for the first time on a gold

SERS substrate, showing the potential of this GNA as a convenient and powerful SERS substrate for

biomolecular detection. In addition, SERS spectrum of R6G on single aggregates has been measured.

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We have shown that the special surface properties of the GNA, in conjunction with strong near IR

absorption, make it useful for SERS analysis of a wide variety of molecules.

KEYWORDS: SERS, Rhodamine 6G, Gold aggregate, nanomaterials, nanoparticle aggregate.

INTRODUCTION

With the discovery of surface enhanced Raman scattering (SERS), the potential for trace molecular

detection using Raman scattering (RS) grew exponentially.¹⁻³ Previously RS had, to a large extent, been

used exclusively for molecular structure interrogations.⁴⁻⁶ Intrinsically small RS signal significantly

limited its practical application. SERS possesses the molecular specificity of Raman scattering with

greatly increased signal, making it ideal for many applications, from trace chemical detection of

biomolecules and explosives⁷⁻¹⁰ to structural interrogations of living cells. 11

The ability to induce SERS is dependent on two key factors: resonant surface plasmon excitation of a

metal substrate and close proximity of analyte molecules to the metal substrate surface. Surface

plasmons are the result of the collective excitation of conduction band electrons near coinage metal

surfaces, namely gold and silver. When a surface plasmon is excited, an enhanced evanescent

electromagnetic (EM) field is created at the metal surface. Calculations have shown that at dislocations

or sharp discontinuities on a metal surface, the EM field will be greatly enhanced. 12,13 Therefore,

molecules in close proximity to the surface will experience a much larger EM field than if they were

directly illuminated with light. The enhanced EM field decays exponentially from the surface and will

induce strong RS of molecules within 0-4 nm of the surface.¹⁴ Further discussion of the SERS

mechanism is left to the many excellent review articles on this subject. 15-18

Significant SERS enhancement is often present at the junction of two or more silver nanoparticles as

shown recently by Brus and co-workers. 19 Little enhancement was observed for single nanoparticles in

this study, however, aggregates with multiple particles yielded large enhancements due to what is

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believed to be the enormous EM field at junction sites. As two metal nanoparticles are brought together to form aggregates, their transition dipoles couple to each other. The enhanced fields of each particle begin to coherently interfere at the junction site between the particles. Calculations by Xu et al have shown that electromagnetic enhancements of 10¹⁰ are present between two nanospheres separated by 1 nm.²⁰ These results suggest that aggregates are better substrates for SERS applications than individual nanoparticles because large enhancements can be achieved at particle junctions of aggregates.²¹ In addition to the works by Brus and Xu, several studies have shown that intentional aggregation of gold and silver nanoparticles significantly increases the observed SERS enhancement.²²⁻²⁷

In addition to enhancing SERS activity of a substrate, aggregation of gold nanoparticles induces plasmon resonances in the near IR (NIR). NIR absorption is highly desirable for many SERS applications, perhaps most notably are in vivo biological studies. Because tissues have an absorption minimum in the NIR, SERS of these materials can best be performed in this spectral region, however, a substrate with NIR absorption is required. Few substrates utilized have been capable of this, however, aggregated gold nanoparticles have strong NIR absorption.²⁸ The development a of SERS substrate such as this is potentially important for detecting biomolecules such as glucose non-invasively, necessary for many biomedical applications.⁷ Not only is there a strong NIR absorption in aggregated gold nanoparticles, but this absorption is tunable by varying particle size and shape.

Because excitation must be on resonance with a maximum number of particles to achieve the largest enhancement possible, the ability to tune the NIR absorption of a SERS substrate is of the utmost importance. The nature of the surface plasmon absorption is entirely dependent on the metal used and the shape of the structure. By changing the structure it is possible to shift the surface plasmon of a metal nanostructure. A spherical nanoparticle of silver, for example, will show a single absorption band near 400 nm, however, if this particle is elongated into a nanorod, a second absorption band between 500 nm and 800 nm will appear depending on length.²⁹ Similarly, in metal nanoshells,³⁰ aggregated nanoparticles,^{31,32} and non-spherical nanoparticles,³³⁻³⁶ new resonances are introduced leading to a variety of extended plasmon bands. For instance, three spherical particles closely interacting in a linear

geometry will have two absorptions similar to a rod shaped particle of the same aspect ratio.³⁷ However, the same particles in a triangular geometry will have more than two resonances.³¹ Therefore, by changing the structure of the nanoshell, nanorod, or aggregate it is possible to effectively tune the position of the plasmon absorption band. This is ideal for SERS as the substrate is tuned to the desired excitation, not vice versa. Aggregates,²³ nanorods,^{24,38} and metal shells³⁹⁻⁴² have been utilized for SERS studies, however, while it is possible to tune the absorption of these structures, aggregates have the advantage of greater enhancements due to intraparticle juncitons.

While aggregates of nanoparticles may lead to the largest SERS enhancements, if the surface is not amicable to the target molecule SERS will not be possible. Surface chemistry of the substrate is critical for SERS since the analyte molecule must be within 0-4 nm of the substrate surface for the electromagnetic field to have an affect. Effectively, the molecule must bind to the metal surface to achieve SERS enhancement, a non-trivial problem for many molecules. For example, rhodamine 6G (R6G), a common dye used in colloidal silver SERS studies, generally does not adsorb to gold surfaces. Such binding selectivity makes engineering a generic SERS substrate challenging. An example of an ideal SERS substrate might be an aggregated nanoparticle system with strong and tunable NIR absorption, and surface properties suitable for binding a variety of chemical and biological analytes. This is the first step to the development of a general analytical technique based on SERS that offers a unique combination of extremely high sensitivity and molecular specificity.

We have recently found that the broad near IR absorption band of a unique gold aggregate system is inhomogeneously broadened due to different sized/shaped aggregates, based on oscillations observed in femtosecond transient absorption and persistent spectral hole-burning experiments. The persistent hole-burning experiment in particular suggests that it may be possible to optically narrow the size/shape distribution of the aggregates for applications such as SERS. This aggregate system has been demonstrated to be SERS active in an early preliminary study.

In this paper we report a detailed study of the SERS activity of this gold nanoparticle aggregate (GNA) system that has unique surface properties and desired near IR absorption for SERS applications.

Utilizing R6G, an ensemble averaged Raman enhancement of approximately 10^7 - 10^9 has been estimated. The strong NIR absorption band, which is tunable between 600-1000 nm, is ideal for biological and physiological studies. NIR Raman also avoids resonance absorption of common biological molecules thereby reducing unwanted background fluorescence. Surface sulfur species allow the easy binding and detection of a variety of molecules, ranging from R6G to several biological molecules never before reported on gold SERS substrates. These molecules have been used to demonstrate the feasibility and versatility of this SERS substrate. The uniquely strong binding character is thought to be due to the high binding affinity of many molecules to the sulfur species on the surface of the GNAs.⁴⁵

EXPERIMENTAL

The synthesis of the gold nanoparticle aggregates has been reported previously 43,46 . Briefly, $500 \,\mu$ l of a $0.02 \,M$ HAuCl₄ stock solution was diluted to $5x10^{-4} \,M$ with Milli-Q water in glassware cleaned in aquaregia and rinsed with Milli-Q water to avoid contamination. To this, $40 - 60 \,\mu$ l of a $0.1 \,M$ solution of Na₂S aged 2-3 months was added. After approximately 60- $120 \,M$ minutes, the color changed from a straw yellow to deep purple with the extended plasmon band (EPB) growing in between 600-1000 nm, indicating reaction completion. The aggregate formation is signified by strong NIR absorption at wavelengths longer than 600 nm. Recent work in our lab has revealed that as the reaction progresses the aggregate structure changes from globular to linear as the EPB shifts from red to blue. The aggregates are stable for weeks or months. After several days some aggregates do settle, but can be dispersed easily with mixing. Absorption measurements were performed on a HP 89532A spectrometer with 2 nm resolution. Raman experiments were performed on a Nicolet Almega Dispersive Raman Spectrometer with excitation at 785 nm, 20 mW of power, and in integration time of 5 seconds averaged over 40 scans.

Samples for single particle experiments were prepared by immobilizing the GNAs on glass coverslips with Trimethoxy[3-(methylamino)propyl]silane (APS). Coverslips were cleaned prior to the silanization

step by sonication in 2% solution of Hellmanex, followed by 18 M Ω water. They were then submerged in 5 mM aqueous solution of APS to deposit the tethering molecules. After 1-2 minutes the coverslips were rinsed with water, dried under nitrogen, and 40 μ L of the aggregate solution was placed on one surface. After several seconds exposed to the solution, it was blown dry with nitrogen.

Single particle experiments were performed on a custom designed confocal microscope built onto an inverted fluorescence microscope (Zeiss, Axiovert 100). A helium-neon laser (633 nm, Melles Griot) was coupled into the back port of the microscope and directed into a high numerical aperture objective (Zeiss Apochromat 100X, 1.4NA) which focused the light onto the sample surface. The sample was then raster scanned across the focused laser to generate an image using a commercially available piezoelectric scanner (Physik Instrumente) and control electronics (Digital Instruments). The Raman scattered light was collected with the same objective used for excitation and focused onto a confocal aperture. The Rayleigh scattered light was then removed using a holographic notch filter (Kaiser Optical) and the remaining Raman scattered light was focused onto an avalanche photodiode (EG&G). Once a nanoparticle aggregate was located, it was centered on the focused laser and the Raman scattering was directed into a spectrograph (Acton Instruments) which dispersed the light onto a liquid nitrogen cooled CCD camera (Princeton Instruments). Typically six spectra (30 seconds) were averaged.

RESULTS AND DISCUSSION

Figure 1 shows a typical electronic absorption spectrum of the gold nanoparticle aggregates. Two plasmon bands are present, corresponding to a transverse plasmon band (530 nm), and an extended plasmon band (EPB) (780 nm). Spherical gold particles will show only the transverse plasmon band due to their totally symmetric geometry. Throughout the reaction, aggregates, not spherical particles, are formed as evidenced by the growth of the EPB indicating strong interparticle interactions. Previous work has shown that particle-particle separation is small, bound together by sulfur species on the nanoparticle surface. While similar absorption characteristics are observed in core/shell and

nanorod structures, it was shown previously that the spectral characteristics here are due to strong particle-particle interation. Interparticle distance is an important factor for SERS studies due to its direct correlation to electromagnetic enhancement. Xu et al have reported that for two 90 nm nanoparticles, decreasing the spacing from 5.5 nm to 1 nm increases the theoretical enhancement by four orders of magnitude. Particles such as these will have greater enhancements than other, weakly interacting systems with less intense EPBs.

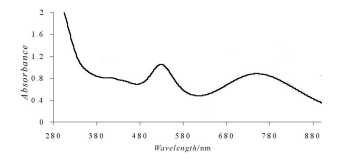


Figure 1. Representative UV-visible spectrum of gold nanoparticle aggregates. The two absorption bands correspond to transverse (530 nm) and extended plasmon bands (peaked around 760 nm).

With slight variation in reagent concentration, it is possible to shift the maximum of the EPB substantially. At standard sodium sulfide concentrations (0.1 mM) the absorption maximum will lie at approximately 780 nm. By increasing the initial sodium sulfide concentration the EPB will blue shift, and at lower concentrations it will shift to the red (figure 2). Three main parameters affect the EPB position: aggregate size, shape, and individual particle size.³¹ Altering any of these factors or combinations thereof will result in drastic changes in peak intensity and position. Generally, as GNAs grow larger, the EPB will red shift. However, aggregate shape also plays an important role in band position.³⁷ Reagent concentration undoubtedly affects not only shape but also size of both the individual nanoparticles and their aggregates, making an experimental distinction between size and shape difficult. The width of the EPB, however, is simply dependent on the polydispersity of the size and shape of the aggregates.^{32,43,50,51}

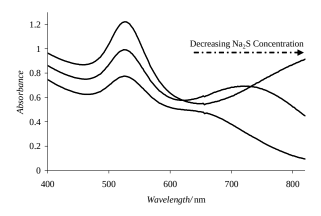


Figure 2. UV-visible absorption spectra of GNAs synthesized with varying concentrations of sodium sulfide. Initial Na₂S concentrations range from 0.8×10^{-4} M to 1.2×10^{-4} M red shifting with a decrease in this concentration.

Because of the random nature of aggregate formation, the GNAs synthesized have a broad distribution of sizes and shapes. Each aggregate with a particular size and shape will have a unique EPB in terms of bandwidth and peak position. However, in an ensemble averaged solution these bands will combine to form one strong, inhomogeneously broadened band. 32,43,50,51 This variety of sizes and shapes is apparent in the transmission electron microscopy (TEM) image in figure 3. Figure 3 is a representative sampling of many different images and samples. The average particle size is on the order of ~20 nm, with the particles making up aggregates, from small to large of many different structures. Each aggregate should have a different EPB depending on its size and shape. 31,32,43,50,51 Because the broad NIR EPB is composed of many narrower absorption bands, upon 785 nm excitation as used in the SERS experiments, only a subset of the aggregates will be on resonance and absorb this incident light. The percentage of aggregates being excited can be maximized by tuning the absorption peak of the broad EPB to be on resonance with the excitation laser. 46 However, the majority of the particles are not excited at a particular excitation wavelength. The exact percentage of aggregates on resonance with a given excitation source is difficult to calculate without knowledge of the homogeneous linewidth of the individual aggregates. To date, we have not been able to determine the homogenous linewidth,

however, it is estimated that as few as 1% of the aggregates are on resonance with a particular excitation wavelength.¹⁹ This parameter is important in estimating the SERS enhancement factor.

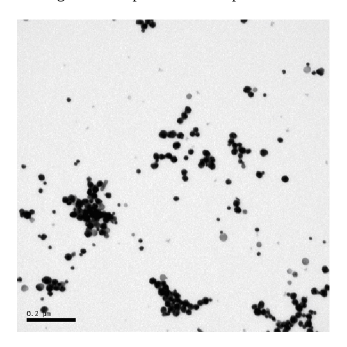


Figure 3. Representative TEM of gold nanoparticle aggregates showing strong interparticle interaction and a variety of aggregate sizes and shapes.

To perform SERS experiments, the aggregate solution EPB maximum was tuned to 785 nm to maximize the number of aggregates on resonance with the excitation laser. It is assumed that the maximum of this band represents the average sized/shaped aggregates. At this excitation wavelength, 785 nm, Raman spectra were first taken without the addition of any analyte. This blank scan, shown in figure 4, shows four strong peaks close to the Rayleigh line at 122 cm⁻¹, 162 cm⁻¹, 195 cm⁻¹, and 258 cm⁻¹. It is assumed that these bands are due to metal-sulfur and/or metal-chloride stretching modes from the adsorbed or chemisorbed sulfur species and chloride ions on the surface, however, the complete assignment is prohibitively difficult at this time due to the complexity of the surface chemistry. Upon addition of any analyte molecule such as R6G, there is a substantial shift in these bands. This is apparent in the disappearance of the 122 cm⁻¹ band and the appearance of the 341 cm⁻¹ band. One possible explanation for this particular shift is the replacement of surface bound chloride with an analyte molecule. The 122 cm⁻¹ peak is likely due to a gold-chloride vibrational mode.⁵² If a surface bound

molecule such as R6G. When the chloride is replaced, the band will shift to a new frequency due to gold-analyte interaction. The exact nature of this mode is unclear athis point. Other low frequency modes are most likely due to gold-sulfur on the surface. Assuming analyte molecules bind to these surface sulfur species, one could possibly explain the small change in SERS spectrum of R6G compared to its normal Raman spectrum by arguing that the direct interaction between R6G and the gold substrate is not strong.

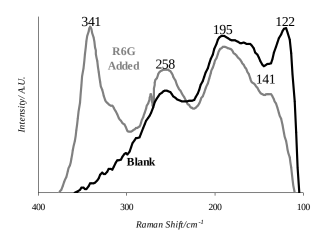


Figure 4. Low energy bands of GNAs alone (blank) and the shifted bands of GNAs with R6G added. The EPB has been tuned to be on resonance with 785 nm excitation.

By simple addition of R6G to the GNA solution, immediate and intense SERS signals were detected with 785 nm excitation (figure 5). It is interesting to note the SERS peaks from R6G are barely shifted, if at all, from the normal Raman peaks of R6G without enhancement. Usually, substantial changes in intensity and shifts in frequency in SERS spectrum can be observed as compared to the normal Raman spectrum of a molecule due to interaction between the molecule and the SERS substrate. The insignificant spectral change observed here may be due to weak interaction between R6G and the SERS substrate. Our observation of little change of SERS spectrum compared to the normal Raman spectrum is similar to previous SERS results of R6G on silver substrates.⁵³ Few studies have shown SERS of R6G from gold substrates because R6G usually does not strongly interact with gold surfaces.^{54,55} In contrast, R6G readily adsorbs to silver and is detectable at nanomolar concentrations and below. The sensitivity of R6G detection by silver allowed the first SERS detection of single molecules.^{21,56} Because

gold substrates have historically been less sensitive to analytes such as R6G, they have been used less frequently. However, there are advantages of gold systems over silver. Silver is easily oxidized while gold is resistant to oxidation and is significantly more stable over time. In addition, gold aggregates can have absorption bands red shifted beyond those of silver, as far as 1 µm, providing more tunability and flexibility in the selection of excitation wavelengths. Core/shell structured substrates can provide similar tunability. However, core/shell structures have so far only exhibited low SERS enhancement factors, because they are not aggregated and lack the concentration of active surface sites of aggregates. In comparison, GNAs afford both tunability and large enhancement values, making aggregates attractive for molecular detection and structural analysis based on SERS. Gold nanoparticles made through the standard citrate reduction can be aggregated to show similar optical properties, however, the surface properties will be substantially different than the GNAs reported here.

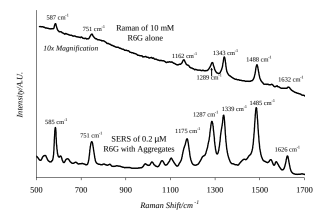


Figure 5. SERS spectrum of 0.2 µM R6G on gold nanoparticle aggregates and Raman spectrum of 10 mM R6G alone (magnified 10x) taken at the same excitation power and collection time. The EPB has been tuned to be on resonance with 785 nm excitation.

The unique surface properties of these GNAs is one of their most important features.⁵⁵ Electron energy loss spectroscopy (EELS) has confirmed the presence of sulfur on the surface of the particles that appears to be effectively capping and linking the aggregates. EELS provides elemental, but not oxidation state information about the surface sulfur. Therefore, the exact oxidation state and molecular

form of these sulfur species are currently unknown and require further study in the future. However, preliminary near edge X-ray absorption fine structure (NEXAFS) measurements have shown that the sulfur species contain sulfur-oxygen bonds, possibly indicating sulfur oxides or simple sulfate ions.⁴⁹ The exact nature of these species is not yet known due to the difficulty of determining the chemical nature of species on the surface of nano-materials. Regardless of their exact oxidation state or molecular form, however, the sulfur species are likely electron rich and are possibly strongly binding towards molecules such as R6G which is positively charged, besides acting as the particle-particle "linker". In addition, the coverage of the surface sulfur species must be very thin, less than 4 nm, making the large SERS enhancements observed possible. The presence of sulfur species makes the surface of these nanoparticles and aggregates unique in comparison to other SERS substrates currently in use.⁵⁷

Table 1. Experimental (exp.) and literature (lit.) SERS shifts of various biologically relevent molecules detected with GNAs^a

Cysteine		<u>Histidine</u>		<u>Lysine</u>		<u>Adenine</u>	
Lit. ⁵⁵	Exp.	Lit. ^{55, 56}	Exp.	Lit. ⁵⁵	Exp.	Lit.54	Exp.
671	671	-	648	620	615	730	735
720	739	801	810	-	672	1339	1320
930	912	931	931	722	741	1482	1463
1067	1057	-	1008	-	868	1581	1564
1140	1136	1160	1134	957	975		
1294	1294	1200	1191	-	1014		
1344	1346	1263	1252	1142	1134		
1411	1394	1312	1322	-	1242		
		1360	1363	1322	1327		
		1448	1463	-	1360		
		1571	1590	1410	1396		
				1464	1467		

<u>Uridine</u>			Tyrosine		<u>Methionine</u>		Leucine	
Lit.	Exp.	Lit. ⁵⁵	Exp.	Lit. ⁵⁵	Exp.	Lit. ⁵⁵	Exp.	
-	698	721	696	700	688	620	642	
-	876	829	810	799	785	718	740	
-	978	-	1053	874	874	-	1240	
-	1016	1199	1217	-	1033	-	1319	
-	1244	1326	1303	1072	1066	1342	1362	
-	1396	1410	1402	1151	1145	1455	1468	
		1613	1593	1353	1350			
		-	1678	-	1390			
				1410	1417			

^aAll positions in units of cm⁻¹

To examine the viability of these GNAs as a general SERS substrate for a variety of applications, several biologically important molecules were tested including adenine, uridine, and several amino

acids. 58-64 Upon the addition of aqueous solutions of these molecules, SERS signals were readily obtained with no further preparation (figures 6 & 7). These spectra compare well with previously reported data as shown in table 1. It is clear that many of the observed peaks were not previously observed in SERS studies. There are two possible explanations for this inconsistency. First, these experiments were all performed at the native pH of the GNA solutions without alteration, i.e. approximately pH 3. As most of the previous work was performed at near neutral pH, this could affect the spectra substantially, changing surface binding and/or protonation of the analyte molecules. As orientation on the surface and molecular structure change with pH, there can be significant changes in the SERS spectrum. Second, the surface of this GNA substrate is very different from any previously studied. The molecules may interact differently with this unique substrate, thereby changing the SERS spectra. In particular, because SERS spectra are dependent on molecular surface orientation, this interaction could have a substantial effect on the observed spectra. A comprehensive discussion of binding orientation and mechanism is beyond the scope of this paper.

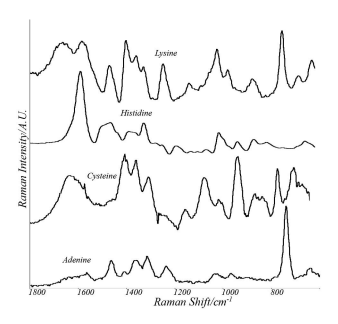


Figure 6. SERS spectra of adenine, cysteine, lysine and histidine in the presence of gold nanoparticle aggregates at pH 3. The EPB has been tuned to be on resonance with 785 nm excitation.

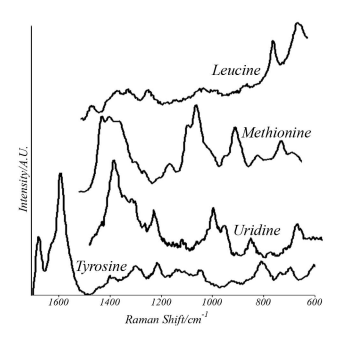


Figure 7. SERS spectra of tyrosine, uridine, methionine, and leucine in the presence of gold nanoparticle aggregates at pH 3. The EPB has been tuned to be on resonance with 785 nm excitation.

While not the first SERS detection of these compounds with the possible exception of uridine, this is their first observation utilizing gold SERS substrates. As mentioned earlier, gold substrates have many advantages over silver, however, until now surface chemistry has hindered its use. With an enhanced SERS-active surface, these GNAs have a distinct advantage over other gold SERS substrates. To provide a semi-quantitative comparison, we will give a rough estimate of the SERS enhancement factor in the current study. Comparing the intensities of the 1500 cm⁻¹ peak of the Raman spectra of 10 mM R6G alone and 0.2 µM R6G in the presence of the aggregates, and including concentration differences, we obtain a 10⁷ enhancement of the Raman signal (figure 5). This enhancement factor is excellent for ensemble averaged, non-resonance SERS in comparison to previous enhancement factors.⁶⁵ If we assume that only about 1% of the aggregates are actually on resonance with the excitation wavelength, the number of R6G molecules involved is reduced by a factor of about 100. 99% of the aggregates with R6G on their surface are not excited and thereby do not contribute to the SERS signal. Taking this factor into account, the enhancement would increase to about 10⁹, which is one of the largest ensemble average enhancement factors reported for SERS substrates.^{24,65,66}

There are two possible reasons for these large enhancements. First, the aggregates have an increased number of SERS active sites due to the great number of nanoparticle junctions present. Because R6G concentration is greater than one molecule per particle in this study, it is possible to have more than one molecule per aggregated enhanced. This also increases the probability of the analyte molecule to adsorb onto an enhancement site or hot spot as opposed to an adsorption site that has no enhanced EM field. Second, the unique surface properties of the GNAs facilitate efficient adsorption or binding of the analyte molecules onto the GNA surface.

To independently confirm the SERS activity of the GNAs, we have measured SERS spectra of R6G on single GNAs using a different experimental setup in a different lab. Utilizing a confocal micro-Raman setup, we are able to measure SERS spectra of individual GNAs. Before applying any analyte solution to the sample slides, blank spectra were taken with no Raman signal observed for any of the examined single GNAs. Upon addition of a solution of R6G to the surface of the sample slide, signal was immediately detected from a large number of individual GNAs (figure 8). It is apparent that the single aggregate SERS signal is substantially different from that of the bulk system. This is not uncommon, and similar observation has been made in single silver aggregate studies. 19,67 This spectral change is due in part to the use of aggregated substrates. The surfaces of the different aggregates can be different and therefore the location of each enhanced molecule varies in terms of surface position. This could change the orientation of the molecule with respect to the active site, for example, in a junction or a dislocation in the surface. Because SERS is relative to the molecule's surface orientation, each aggregate can have a slightly different SERS signal, however, the bulk solution will average to the expected spectrum observed in figure 5. Molecules were able to adsorb onto the aggregates in film form to yield strong SERS signals. This is encouraging particularly for applications requiring SERS measured in solid films. One important feature of a practical SERS sensor or detection device would be reusability of the substrate. This is only possible if the molecules detected can be readily removed so the GNA substrate can be reused. This was tested by removing the R6G by rinsing the substrate with water, at which point no SERS spectra were observed. With a fresh drop of R6G applied to the GNA substrate, equally strong SERS signals can be measured again. Therefore, the films of these GNAs are not only stable as mentioned earlier but also reusable, a very useful trait for many potential SERS applications.

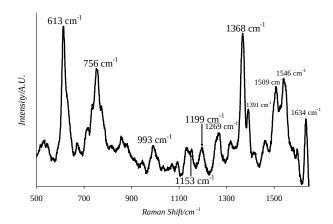


Figure 8. Representative single particle SERS spectrum of R6G on single gold nanoparticle aggregates immobilized to glass coverslips. The EPB of the GNA solution was tuned to maximize the number of resonant particles with excitation at 633 nm.

Conclusion

A unique gold aggregate system has been shown experimentally to be useful for SERS detection of R6G and several biological molecules. R6G, a common molecule for SERS studies that is generally difficult to detect on gold substrates, readily adsorbs to these GNAs. This is believed to be due to the presence of sulfur species on the particle surface that provide strong binding of molecules of interest to the GNA. The SERS activity demonstrated for adenine, cysteine, lysine, histidine, leucine, methionine, uridine and tyrosine shows the potential of the GNAs as convenient SERS substrates for biological and chemical sensing and analysis. Based on the surface properties of these GNAs, a wide variety of molecules can be detected via this unique SERS substrate which has been shown to be active for many molecules never before detected on gold substrates. These GNAs were shown to yield a large SERS enhancement of 10⁷-10⁹ for R6G. The large enhancement is partly attributed to aggregation that has been shown to substantially increase SERS signals. The potential of SERS using GNA films was

tested based on single aggregate studies. It was found that R6G was immediately detectable with individual single aggregates immobilized onto glass slides, and the substrate appears to be reusable.

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